

In the Claims:

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- 1. (currently amended) A method for processing dermal tissue for implantation into a subject, said method comprising the steps of:
- a. removing the epidermal layer of said dermal tissue to produce de-epidermalized tissue;
- b. incubating said de-epidermalized tissue in at least one processing solution to remove cells from said de-epidermalized tissue, thereby producing a decellularized tissue matrix;
- c. treating said decellularized tissue matrix to cause a reduction in size and an increase in surface area; and
- e. d. exposing said decellularized tissue matrix to an acylating agent, wherein the ratio of said acylating agent to wet tissue-weight is about 0.1% to about 0.3% 0.003:1 or less, thereby producing a dispersed tissue matrix.
- Canceled (incorporated into claim 1).
- 3. (original) The method of claim 2, wherein said treating comprises cryomilling said decellularized tissue matrix.
- 4. (original) The method of claim 1, further comprising contacting said de-epidermalized tissue with a viral inactivating agent, before, after, or during step (b).
- 5. (original) The method of claim 1, wherein said tissue is mammalian.
- 6. (original) The method of claim 4, wherein said tissue is human.
- 7. (original) The method of claim 1, wherein said acylating agent is glutaric anhydride or succinic anhydride.



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- 8. (currently amended) The method of claim 1, wherein said ratio of acylating agent-towet-tissue weight is about 0.002:1 to about 0.001:1 0.1% to about 0.2%!
- 9. (original) The method of claim 1, wherein said decellularization solution comprises sodium hydroxide.
- 10. (original) The method of claim 1, wherein said decellularization solution comprises phosphoric acid.
- 11. (original) The method of claim 1, wherein said tissue is autogenic, allogenic or xenogenic.
- 12. (original) The method of claim 1, wherein said step of removing the epidermal layer comprises exposing said tissue to a hypertonic salt solution.
- 13. (currently amended)A method for dispersing decellularized animal <u>connective</u> tissue, said method comprising <u>the steps of</u>:

treating said decellularized animal connective tissue to cause a reduction in size and an increase in surface area; and

contacting said decellularized, treated, animal connective tissue with a solution comprising an acylating agent, wherein the ratio of said acylating agent to wet tissue weight is about 0.1% to about 0.3% 0.003:1 or less.

- 14. Canceled (incorporated into claim 13).
- 15. (original) The method of claim 14, wherein said treating comprises cryomilling said decellularized tissue.
- 16. (original) The method of claim 13, wherein said tissue is mammalian.





- 17. (original) The method of claim 13, wherein said tissue is human.
- 18. (original) The method of claim 13, wherein said tissue is connective tissue.
- 19. (original) The method of claim 13, wherein said tissue is dermal tissue.
- 20. (currently amended) The method of claim 13, wherein said ratio of acylating agent to wet tissue weight is about 0.002:1 to about 0.001:1 0.1% to about 0.2%.
- 21. (currently amended) A method for altering augmenting the condition of in situ tissue of a subject, said method comprising introducing an effective amount of a dispersed collagen matrix being at the site of the into said in situ tissue of said subject, said dispersed collagen matrix being prepared by treating a decellularized animal connective tissue matrix to cause a reduction in size and an increase in surface area and contacting said decellularized, treated animal connective tissue matrix with a solution comprising an acylating agent, wherein the ratio of said acylating agent to wet tissue weight is about 0.1% to about 0.3% 0.003:1 or less.
- 22. (original) The method of claim 19, wherein said subject is a human.
- 23. (original) The method of claim 19, wherein said dispersed collagen matrix is derived from an allogeneic source.
- 24. (original) The method of claim 1, wherein said acylating agent is glutaric anhydride or succinic anhydride.
- 25. (currently amended) The method of claim 1, wherein said ratio of acylating agent to wet tissue weight is about 0.002:1 to about 0.001:1 0.1% to about 0.2%.
- (currently amended) A composition comprising an injectable, dispersed collagen



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matrix prepared by <u>treating a decellularized animal connective tissue matrix to cause a</u> reduction in size and an increase in surface area and contacting <u>said</u> decellularized, <u>treated</u> animal connective tissue with a solution comprising an acylating agent, wherein the ratio of said acylating agent to wet tissue weight is about <u>0.1%</u> to about <u>0.3%</u> 0.003:1 or less.

- 27. (original) The composition of claim 26, wherein the dispersed collagen matrix is injectable through a 30 gauge needle.
- 28. (currently amended) The composition of claim 26, wherein the dispersed collagen matrix has a trypsin resistance such that greater than about 40% of the dispersed collagen matrix remains undigested when exposed to 2% trypsin at 37°C for 6-24 hours.
- 29. (currently amended) The composition of claim 26, wherein the dispersed collagen matrix has a trypsin resistance such that greater than about 50% of the dispersed collagen matrix remains undigested when exposed to 2% trypsin at 37°C for 6-24 hours.
- 30. (currently amended) The composition of claim 27, wherein the dispersed collagen matrix has a trypsin resistance such that greater than about 70% of the dispersed collagen matrix remains undigested when exposed to 2% trypsin at 37°C for 6-24 hours.
- 31. (currently amended) The composition of claim 27, wherein the dispersed collagen matrix has a trypsin resistance such that greater that about 90% of the dispersed collagen matrix remains undigested when exposed to 2% trypsin at 37°C for 6-24 hours.
- Deleted.
- 33. (currently amended) An injectable composition comprising an <u>decellularized</u>, acylated, dispersed, dermal tissue matrix having a trypsin resistance <u>such that</u> greater than about 40% <u>of the dispersed collagen matrix remains undigested when exposed to 2% trypsin at 37°C for 6-24 hours.</u>



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- 34. (currently amended) The composition of claim 33, wherein the dermal tissue matrix has a trypsin resistance such that greater than about 50% of the dispersed collagen matrix remains undigested when exposed to 2% trypsin at 37°C for 6-24 hours.
- 35. (currently amended) The composition of claim 33, wherein the dermal tissue matrix has a trypsin resistance such that greater than about 70% of the dispersed collagen matrix remains undigested when exposed to 2% trypsin at 37°C for 6-24 hours.
- 36. (currently amended) The composition of claim 33, wherein the dermal tissue matrix has a trypsin resistance such that greater than about 90% of the dispersed collagen matrix remains undigested when exposed to 2% trypsin at 37°C for 6-24 hours.